

The effect of temperature on the phenotypic features and the maceration ability of *Dickeya solani* strains isolated in Finland, Israel and Poland

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Abstract Pectinolytic bacteria from the genus *Dickeya* (former *Erwinia chrysanthemi*), belonging to *Dickeya dianthicola* and *Dickeya solani* species, are causative agents of blackleg and soft rot diseases in Europe. Recently, *D. solani* have been isolated most frequently from potato plants with the symptoms of blackleg and soft rot. *D. solani* strains were shown to cause more severe disease symptoms on potato plants than *D. dianthicola* especially at the higher temperature. They are also able to develop blackleg disease from lower inoculum levels. In the presented study we not only compared phenotypic features of fifteen *D. solani* strains isolated in countries having different climatic conditions, Poland, Finland and Israel, but also we examined three *D. dianthicola* strains. The comparison was performed to determine the influence of the strain origin and the temperature of incubation on the ability of the strains to macerate potato tissue and on their major virulence factors such as: pectinolytic, cellulolytic and proteolytic activities, siderophore production and motility. Polish *D. solani* strains showed higher activities of cell wall degrading enzymes than the Finnish and Israeli strains at all the tested temperatures: 18, 27, 37 °C. This

observation is correlated with the higher ability of Polish *D. solani* strains to cause soft rot. In addition, *D. solani* strains exhibited higher activity of the above mentioned enzymes and caused more severe potato tuber maceration in laboratory tests than the tested *D. dianthicola* strains. The collected results indicate that although *D. solani* strains from different climatic conditions have identical Pulse Field Gel Electrophoresis (PFGE) profiles in addition to the same fingerprint profiles obtained by the repetitive sequence-based polymerase chain reaction (REP, ERIC and BOX repetitive sequences), they differ in the examined phenotypic features, especially in the activities of pectinolytic, cellulolytic and proteolytic enzymes and their capacity to macerate potato tuber tissue.

Keywords Different climate · Pectinolytic bacteria · Plant cell-wall degrading enzymes · Rep-PCR · PFGE · Virulence factors

Introduction

Dickeya species (Samson et al. 2005), formerly named *Pectobacterium chrysanthemi* (Hauben et al. 1998) and *Erwinia chrysanthemi* (Burkholder et al. 1953), are causative agents of soft rot and blackleg in potato. Similarly to *Dickeya* species, *Pectobacterium* species, formerly called *Erwinia carotovora* (Burkholder et al. 1953), can cause blackleg and soft rot. The genus *Dickeya* is a highly diverse group, which according to the present classification contains eight species: *D. aquatica*, *D. chrysanthemi*, *D. dadantii*, *D.*

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dianthicola, *D. fangzhongdai*, *D. paradisiaca*, *D. solani* and *D. zae* (Samson et al. 2005; Brady et al. 2012; van der Wolf et al. 2014; Parkinson et al. 2014; Tian et al. 2016). Bacteria from this genus were known to cause disease symptoms, especially in tropical and subtropical climates, on a large number of host plant species, including many economically important crops like potato, maize and banana (Ma et al. 2007). *Dickeya* sp. strains isolated from potato plants outside of Europe fell into biovar 3 and 6 and possessed high optimum growth temperature. They were classified to *D. dadanti*, *D. chrysanthemi* or *D. zae* species (Dickey 1981; Samson et al. 2005; Toth et al. 2011).

Dickeya strains that were firstly isolated from potato plants in Europe belonged to biovar 1 and 7, which coincide with *D. dianthicola*, a species adapted to cool regions (Cazelles and Schwarzel 1992; Janse and Ruissen 1988). During last decade, strains from *D. solani* species were isolated from potatoes across Europe and Israel (Toth et al. 2011; van der Wolf et al. 2014) undoubtedly more often than the strains of *D. dianthicola*. Laurila et al. (Laurila et al. 2008; Laurila et al. 2010) and Degefu et al. (2013) reported the presence of *D. solani* strains in Finland. Also, the strains of *D. solani* were isolated from potato plants with blackleg and soft rot symptoms in France, the Netherlands, Poland, Spain and Switzerland (Palacio-Bielsa et al. 2006; Helias 2006; Czajkowski et al. 2009; Slawiak et al. 2009a; b; Toth et al. 2011; Gill et al. 2014; Potrykus et al. 2016). Besides, Tsror et al. (Tsror (Lahkim) et al. 2009; Tsror (Lahkim) et al. 2013) isolated *D. solani* strains from blackleg diseased potato plants in Israel.

In the recent years, potato losses are caused, apart from the *Dickeya* strains, also by other pectinolytic bacteria known as tropical and subtropical pathogens, e.g. *Pectobacterium wasabiae* (Nabhan et al. 2012; Waleron et al. 2013), and *Pectobacterium carotovorum* subsp. *brasiliense* (de Werra et al. 2015; Nunes Leite et al. 2014; Waleron et al. 2015). These losses caused by the strains from the mentioned taxa increased significantly in a large number of European countries. We could hypothesize, that the fast spread of *D. solani*, *P. wasabiae* and *P. c.* subsp. *brasiliense* in Europe is connected with an increase in the spring and summer temperatures, arising as a consequence of the global warming and more intense international seed potato exchange.

Bacteria from species *D. solani* have a wider optimal temperature range for growth and soft rot disease development (from 25 °C to 39 °C). They also possess

features which allow for more effective colonization and faster spread through the plant vascular system than in the case of *D. dianthicola* (Czajkowski et al. 2010). All previously characterized *D. solani* strains indicated identical rep-PCR and PFGE profiles (Degefu et al. 2013; Tsror (Lahkim) et al. 2013; van der Wolf et al. 2014; Potrykus et al. 2016). However, the comparison of genome sequences of several *D. solani* strains indicated the existence of some differences within the studied strains caused probably by the horizontal gene transfer (Khayati et al. 2015).

Crucial factors for soft rot and blackleg symptoms development influencing the potato tissue maceration are, amongst other aspects, the growth rate of the pathogen, its capacity to produce plant cell-wall degrading enzymes (PCWDE) and its motility (Hugouvieux-Cotte-Pattat et al. 1996). The *Dickeya* group is characterized by the ability to produce pectinolytic enzymes, among which pectate lyases are considered as the most important virulence factors. Extracellular pectate lyases cause degradation of pectins and polygalacturonic acid, which are important components of the plant cell wall. Studies on a model strain *D. dadantii* 3937 indicated that the specific regulation of the pectinases expression and secretion depends on the temperature, humidity and the type of plant tissue (Hugouvieux-Cotte-Pattat et al. 1996; Hugouvieux-Cotte-Pattat and Charaoui-Boukerzaza 2009; Hugouvieux-Cotte-Pattat et al. 2014; Kazemi-Pour et al. 2004). According to Hugouvieux-Cotte-Pattat et al. (Hugouvieux-Cotte-Pattat et al. 1996) temperature is one of the most important external factors influencing the survival of *D. dadantii* 3937 in the environment. Temperature also modifies the growth rate and the metabolic activity of this pathogen and its ability to cause disease symptoms. However, little is known about the influence of environmental conditions on pectinolytic enzymes production and the aggressiveness of *D. solani*.

The question why *D. solani* has spread throughout Europe and caused more severe losses than *D. dianthicola* has been raised by researchers across Europe (Czajkowski et al. 2009; Toth et al. 2011; Degefu et al. 2013; Potrykus et al. 2016). Studies show that these pathogens are able to induce diseases at the lower inoculum levels, produce more PCWDE and in addition possess an arsenal of T5SS/T6SS secreted proteins exhibiting more diverse functions than just destroying the plant cell-wall (Toth et al. 2011; Garland et al. 2013; Pedron et al. 2014; Khayati et al. 2015; Potrykus et al. 2016).

Strains of *D. solani* are able to cause severe disease symptoms in a wide range of climates (for example in European countries like Finland, the Netherlands, Poland, France, or Switzerland, to a country such as Israel). Yet, there is only one comparative study performed on *D. solani* strains originating from different climate conditions (Tsrer (Lahkim) et al. 2013). In the present study, we verified the genetic homogeneity of the strains of *D. solani* isolated in the previous years from infected potato plants under different temperature conditions (in Poland, Finland and Israel) and compared their phenotypic features such as pectinases, cellulases and proteases activities, siderophores production, potato tissue maceration, as well as their motility. The main goal of this study was to evaluate the effect of the origin of isolation (Finland, Israel, Poland), the

incubation temperature (18, 28 and 37 °C) on the intensiveness of the tested features. The above shown temperatures were arbitrary chosen as typical for the summer months in cold, temperate and hot climate countries.

Materials and methods

Bacterial strains, media and culture conditions

D. solani strains isolated in Finland, Israel and Poland and strains from different *Dickeya* species used in this paper are listed in Table 1. Polish *Dickeya solani* isolates were obtained in 2005, 2009 and 2010 during routine procedures of isolating bacteria from infected seed potato

Table 1 *Dickeya* spp. strains used in this study

Name	IFB number (other collections)	Country and year of isolation	Source, references
<i>Dickeya dadantii</i>	IFB0010 ^{a,b} (NCPBP898 ^T , IPO2120)	Comoros, 1960	<i>Pelargonium capitatum</i> Samson et al. 2005
<i>Dickeya dadantii</i>	IFB0016 ^c (3937, A4922)	France	<i>Sepolia ionantha</i> Kotoujansky et al. 1982
<i>Dickeya dianthicola</i>	IFB0103 ^a (NCPBP453 ^T , IPO2114)	UK, 1956	<i>Dianthus caryophyllus</i> Samson et al. 2005
<i>Dickeya dianthicola</i>	IFB0157 ^d (LMG28825)	Poland, 2009	<i>Solanum tuberosum</i> This work
<i>Dickeya dianthicola</i>	IFB0188 ^a (IPO1741)	The Netherlands, 1992	<i>Solanum tuberosum</i> van der Wolf et al. 2014
<i>Dickeya solani</i>	IFB0123 ^a (IPO2222)	The Netherlands, 2007	<i>Solanum tuberosum</i> Czajkowski et al. 2013
<i>Dickeya solani</i>	IFB0099 ^d (IPO2276, LMG28824)	Poland, 2005	<i>Solanum tuberosum</i> Slawiak et al. 2009a, b
<i>Dickeya solani</i>	IFB0100 ^d (IPO2277)	Poland, 2005	<i>Solanum tuberosum</i> This work
<i>Dickeya solani</i>	IFB0158 ^d (LMG 28826)	Poland, North 2009	<i>Solanum tuberosum</i> Potrykus et al. 2014
<i>Dickeya solani</i>	IFB0167 ^d	Poland, South-west 2009	<i>Solanum tuberosum</i> Potrykus et al. 2016
<i>Dickeya solani</i>	IFB0212 ^d	Poland, Central 2010	<i>Solanum tuberosum</i> This work
<i>Dickeya solani</i>	IFB0231 ^e (VIC-BL 25)	Finland 2008	<i>Solanum tuberosum</i> Degefu et al. 2013
<i>Dickeya solani</i>	IFB0236 ^e (VIC-SRI 5)	Finland 2008	<i>Solanum tuberosum</i> Degefu et al. 2013
<i>Dickeya solani</i>	IFB0254 ^e (SRG 5–4)	Finland 2008	<i>Solanum tuberosum</i> Degefu et al. 2013
<i>Dickeya solani</i>	IFB0261 ^e (BLG 4–5)	Finland 2008	<i>Solanum tuberosum</i> Degefu et al. 2013
<i>Dickeya solani</i>	IFB0265 ^e (SRG 54–4)	Finland 2008	<i>Solanum tuberosum</i> Degefu et al. 2013
<i>Dickeya solani</i>	IFB0124 ^f (IPO3228)	Israel, 2008	<i>Solanum tuberosum</i> Tsrer (Lahkim) et al. 2013
<i>Dickeya solani</i>	IFB0125 ^f (IPO3296)	Israel, 2008	<i>Solanum tuberosum</i> Tsrer (Lahkim) et al. 2013
<i>Dickeya solani</i>	IFB0455 ^f (IPO3204)	Israel, 2007	<i>Solanum tuberosum</i> Tsrer (Lahkim) et al. 2013
<i>Dickeya solani</i>	IFB0456 ^f (IPO3213)	Israel, 2008	<i>Solanum tuberosum</i> Tsrer (Lahkim) et al. 2013
<i>Dickeya solani</i>	IFB0457 ^f (IPO3236)	Israel, 2009	<i>Solanum tuberosum</i> Tsrer (Lahkim) et al. 2013

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^f Leah Tsrer, Israel

as reported by Potrykus et al. (2016). Five *D. solani* strains isolated in Finland were obtained by courtesy of Dr. Yeshitila Degefu, MTT Agrifood Research Finland, Biotechnology and Food Research, Agro Biotechnology University of Oulu, FI, Finland. Israeli strains were obtained by courtesy of Dr. Leah Tsrer, Gilat Research Center, Israel (five strains) and Dutch strain (*D. solani* Type strain IPO2222) thanks to Dr. Jan. van der Wolf, Plant Research International, Wageningen.

All strains were stored in 40 % glycerol at -80°C . Bacteria were cultured in LB medium or LB supplemented with agar 15 g L^{-1} and/or CVP (Crystal Violet Pectate, Helias et al. 2012) at 28°C for 24–48 h prior to DNA extraction and performing the phenotypic and maceration ability tests. In the case of liquid preparations, bacterial cultures were grown in LB medium at 28°C with shaking (200 rpm).

Genetic characterization of *D. solani* strains from different climate conditions

REP-PCR profiling

Finish, Israeli and Polish strains of *D. solani* and the Type Strains of other *Dickeya* species (Table 1) were analyzed using the repetitive sequence based PCR (rep-PCR) with three sets of primers: ERIC 1R and ERIC 2 (Enterobacterial Repetitive Intergenic Consensus), REP 1 R-I and REP 2-I (Repetitive Extragenic Palindromic sequences) and also BOX1AIR as described by Versalovic et al. (Versalovic et al. 1991 and Versalovic et al. 1994), respectively. Genomic DNA was isolated using Genomic Mini AX Bacteria Kit (A&A Biotechnology, Poland). The resulting products of amplification with REP, ERIC or BOX primers were separated by electrophoresis at 50 V for 2.5 h in 0.8 % agarose (Basica, Prona) gel using 0.5 x TBE buffer and visualized and documented using the Gel Doc imaging system (Bio Rad Laboratories Inc.) supported by Quantity One Software (Bio Rad Laboratories Inc., Hercules, CA, USA).

Restriction fragment length polymorphism - pulse field gel electrophoresis (RFLP-PFGE)

The RFLP - PFGE was performed as already described by Degefu et al. (2013). After the electrophoresis run was completed, the gels were stained with ethidium bromide solution (0.5 mg mL^{-1}), rinsed with water, and the band pattern was observed under UV light. As a

standard, *Salmonella enterica* serotype Braenderup (strain H9812, ATCC) was used.

D. solani growth curves determination

Determination of the growth dynamics was established in 48-well plates with the use of EnVision spectrophotometer plate reader (Perkin Elmer, USA) with orbital shaking at two temperatures: 28 and 37°C . The night culture of each strain was diluted 1: 50 in fresh LB medium, reaching $500\text{ }\mu\text{L}$ as the final volume of the culture. The measurement of the absorbance at 600 nm was performed every hour for 20 h. The experiment with two replicates was performed twice.

Qualitative phenotypic analyses – plate assays

For plate tests, a bacterial suspension containing 10^8 cfu mL^{-1} was prepared and $2\text{ }\mu\text{L}$ was spotted on a plate and incubated for 48 h, unless otherwise stated. All experiments were performed at three different temperatures: 18 , 28 and 37°C . The experiments with two replicates were performed three times.

Pectinase activity was tested on plates with M63Y PGA medium (Reverchon et al. 1986). After 24 h incubation plates were covered with 10 % copper acetate solution. Copper acetate reacts with PGA, giving the blue color of medium. If bacteria degrade PGA there is a whitish halo around bacterial colonies. The halo diameter is in proportion to the activity of pectinases.

Cellulase activity was analyzed on M63 agar plates supplemented with glycerol at 2 g liter^{-1} and carboxymethylcellulose at 10 g L^{-1} , as described by Wood (1980). After incubation the plates were flooded with Congo Red solution (10 mg mL^{-1}) for 10 min and washed for 5 min with 1 M NaCl . The diameter of the clear halo around the colonies was measured. The halo diameter indicated the activity of cellulases.

Protease activity was measured on medium supplemented with skim milk (12.5 g L^{-1}) (Ji et al. 1987). The diameter of the clear halo around the colonies was measured. The halo diameter is in proportion to the activity of produced proteases.

Siderophore production and an **ability to chelate iron ions** were determined on chrome azurol S-agar plates (Schwyn and Neilands 1987). The diameter of the orange halo around the colonies was measured.

Motility assay. To assess the swimming motility bacterial strains were inoculated onto 0.3 % semisolid

agar plates (Harshey 2003) and incubated for 24 h. The diameter of the colonies was measured.

Quantitative evaluation of pectate lyases activities

For measurements of the total pectate lyase activity, bacteria were grown in M63 medium (Miller 1972) supplemented with glycerol (Y) to the final concentration of 2 % (noninduced conditions) or glycerol and poligalacturonic acid (PGA, Sigma) (induced conditions) to the final concentration of 2 and 0.2 %, respectively. The bacteria were cultivated up to the stationary phase (24 h) in aerobic conditions at 28 °C with shaking (200 rpm).

The assay medium contained PGA 0.05 %, 0.1 mM CaCl_2 , 100 mM Tris- HCl pH 8.5. This pH was chosen as optimal for majority of pectate lyase (Hugouvieux-Cotte-Pattat et al. 2014). Pectate lyase activity (PL) was determined by spectrophotometric monitoring of the ability to form unsaturated products, which absorb light at 235 nm (Tardy et al. 1997). Bacterial cultures were grown to the stationary phase (24 h). One unit of pectate lyases activity was defined as the amount of enzyme required to produce 1 μmol of unsaturated product per 1 min. Total pectate lyases activity was expressed as μmoles of unsaturated products liberated per min per mg of bacterial dry mass. The experiment with three replicates was performed three times.

Maceration ability test

Potatoes cv. Lord, average tuber size 60–70 mm were washed under running water and sterilized with 10 % of commercially available bleach (i.e. ACE or Clorox) for 20 min. Later on, tubers were thoroughly washed in water. Tubers were dried under the laminar flow. Dried tubers were cut aseptically crosswise into 10 mm thick slices. Slices were placed in glass Petri dishes (150 mm \times 25 mm) containing a layer of Whatman filter paper. Depending on the slice size, two or three holes 5 \times 5 mm were cut in the surface of the slices. Into each hole 50 μl of bacterial suspension was added. Whatman filter paper in each Petri dish was moistened with 5 ml of sterile water. Potato slices were inoculated with three different level of bacteria (10^5 , 10^6 and 10^7 cfu ml^{-1}). Bacterial suspensions were prepared in 0.85 % NaCl. Inoculated slices were incubated at 28 °C for 48 h and the diameter of rotten tissue was measured. As a negative control 50 μl of 0.85 % NaCl was used. The experiment with nine replicates was repeated twice.

Statistical analysis

Statistica 12 ® (StatSoft Inc., USA) was used to analyze data related to phenotypic traits of *Dickeya* spp. strains (enzymatic activities, siderophore production and swimming motility in relation to temperature, also pectinolytic activity test and ability to macerate potato tuber slices). For the dataset of enzymatic activities, siderophores production and motility, a two-way analysis of variance (ANOVA) was performed where ‘origin of strain’ or ‘species’ and ‘temperature’ were used as the experimental factors. For the dataset of total pectinolytic activity, a two-way ANOVA was conducted with ‘origin of strain’ or ‘species’ and ‘conditions’ taken as the experimental factors. For the dataset of ability to macerate potato tuber slices the two-way ANOVA was conducted with ‘origin of strain’ and ‘inoculation level’ as the experimental factors. The impacts of the origin/species and the temperature on tested phenotypic traits were derived from the one-way ANOVA models as the determination coefficients (presented in percentages).

Results

Molecular profiling of *D. Solani* strains from different climatic conditions

Two molecular profiling techniques: rep-PCR and RFLP-PFGE were used to characterize the genomic material of *D. solani* strains isolated in Poland, Finland and Israel. The results of fingerprinting indicated that genomic profiles of all tested *D. solani* strains were the same regardless of the profiling method used (Fig. 1). Profiles of *D. solani* from Poland, Finland and Israel have the same molecular pattern as *D. solani* IPO 2222^{TS} (IFB0123). At the same time they do differ from the profiles of *D. dadantii* (strains IFB0010^{TS} and IFB0016) and *D. dianthicola* (strains IFB0103^{TS} and IFB0157).

Dickeya sp. growth dynamic

All the tested *D. solani* strains have a similar growth dynamic. At 28 °C, *D. solani*^{TS} strain reaches the stationary phase after the same time as *D. dadantii*^{TS} (after 12 h), earlier than Polish, Finnish, Israeli strains of *D. solani* and *D. dianthicola*^{TS} (13.8 h, 15.6 h, 12.8 h and 14 h, respectively). At 37 °C *D. dianthicola*^{TS} reaches stationary phase most rapidly (after 12 h), earlier

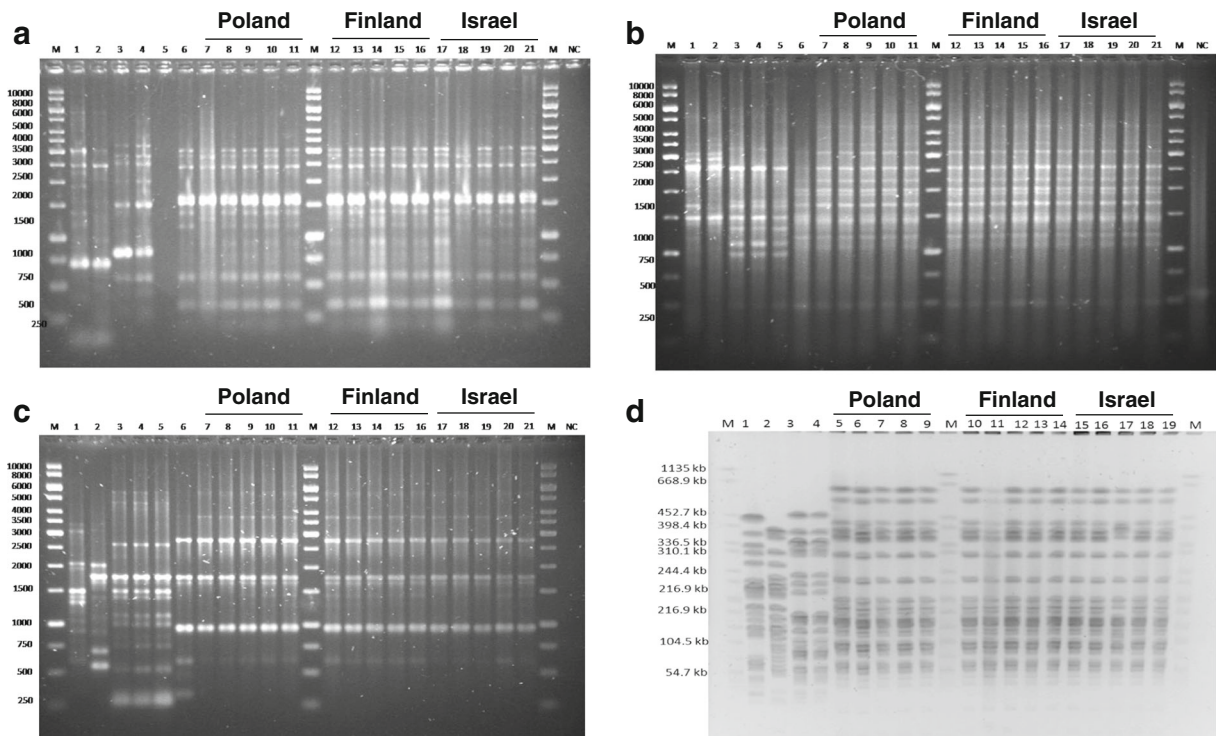


Fig. 1 Genetic homogeneity of tested *D. solani* strains evaluated by rep-PCR and RFLP-PFGE profiling. **a** REP PCR profiling, **b** BOX PCR profiling, **c** ERIC PCR profiling. For panels **a**, **b** and **c** the order is as follows: 1 - *D. dadantii* IFB0010; 2 - *D. dadantii* IFB0016; 3 - *D. dadantii* IFB0016; 4 - *D. dadantii* IFB0016; 5 - *D. dadantii* IFB0016; 6 - *D. dadantii* IFB0016; 7 - *D. dadantii* IFB0016; 8 - *D. dadantii* IFB0016; 9 - *D. dadantii* IFB0016; 10 - *D. dadantii* IFB0016; 11 - *D. dadantii* IFB0016; 12 - *D. dadantii* IFB0016; 13 - *D. dadantii* IFB0016; 14 - *D. dadantii* IFB0016; 15 - *D. dadantii* IFB0016; 16 - *D. dadantii* IFB0016; 17 - *D. dadantii* IFB0016; 18 - *D. dadantii* IFB0016; 19 - *D. dadantii* IFB0016; 20 - *D. dadantii* IFB0016; 21 - *D. dadantii* IFB0016; M - marker; NC - negative control

- IFB0455; 20 - IFB0456; 21 - IFB0457. **d** PFGE profiling, 1 - *D. dadantii* IFB0010; 2 - *D. dadantii* IFB0016; 3 - *D. dadantii* IFB0016; 4 - *D. dadantii* IFB0016; 5 - *D. dadantii* IFB0016; 6 - *D. dadantii* IFB0016; 7 - *D. dadantii* IFB0016; 8 - *D. dadantii* IFB0016; 9 - *D. dadantii* IFB0016; 10 - *D. dadantii* IFB0016; 11 - *D. dadantii* IFB0016; 12 - *D. dadantii* IFB0016; 13 - *D. dadantii* IFB0016; 14 - *D. dadantii* IFB0016; 15 - *D. dadantii* IFB0016; 16 - *D. dadantii* IFB0016; 17 - *D. dadantii* IFB0016; 18 - *D. dadantii* IFB0016; 19 - *D. dadantii* IFB0016; 20 - *D. dadantii* IFB0016; 21 - *D. dadantii* IFB0016; M - marker; NC - negative control

than *D. dadantii* (13 h) and *D. solani*^{TS} (16 h). Interestingly, at 37 °C Polish strains grew to the stationary phase more quickly than the Finnish and Israeli ones: 14.8 h in comparison to 15.6 and 16 h, respectively (Table 2).

The statistical analysis of the time of entering the stationary phase as well as the level of the maximum optical density reached at 28 °C and at 37 °C did not allow to distinguish significant differences in growth dynamics between strains of *D. solani* originating from different climatic conditions, nor between different *Dickeya* species (one-way ANOVA, $p > 0.05$).

Phenotypes of *D. solani* strains originating from different climatic conditions

Strains originating from Poland are the most homogeneous group, within the group of Finnish strains there is

usually one strain exhibiting lower enzymatic activities, while the Israeli strains are the most diverse, especially regarding enzymatic activities and the ability to swim (Supplement 1). All tested *Dickeya* strains have the highest enzymatic activities at 28 °C. Polish *D. solani* strains exhibit the highest pectinolytic, cellulolytic and proteolytic activities at this temperature (Fig. 2). They differ significantly from the Israeli strains in all tested temperatures. The enzymatic activity of Finnish *D. solani* strains comprises between the levels determined for Polish and Israeli strains.

Israeli strains indicated the highest siderophores production (ability to chelate iron ions) at 28 °C, but not at 18 and 37 °C. The Finnish strains did not differ significantly from the Israeli and the Polish ones (Fig. 2). Besides, Polish and Finnish strains indicated significantly better ability to swim at 28 °C, then Israeli strains, but at other temperatures differences were not significant (Fig. 2).

Table 2 Growth of *D. solani* strains at 28 °C and 37 °C in comparison to *D. dadantii* and *D. dianthicola*

<i>Dickeya</i> spp.		Time to achieve maximum density [h]		Maximal optical density	
		28 °C	37 °C	28 °C	37 °C
<i>D. solani</i> from Poland	IFB0099	14	15	0.92	0.86
	IFB0100	15	13	0.90	0.87
	IFB0158	13	14	0.84	0.89
	IFB0167	14	16	0.88	0.77
	IFB0212	13	16	0.82	0.80
	Average	13.8	14.8	0.87	0.84
<i>D. solani</i> from Finland	IFB0231	16	16	0.68	0.85
	IFB0236	16	13	0.64	0.50
	IFB0254	15	16	0.81	0.79
	IFB0261	16	16	0.79	0.85
	IFB0265	15	16	0.82	0.94
	Average	15.6	15.4	0.75	0.78
<i>D. solani</i> from Israel	IFB0124	10	16	0.86	0.71
	IFB0125	16	16	0.73	0.86
	IFB0455	15	16	0.87	0.68
	IFB0456	13	16	0.59	0.51
	IFB0457	10	16	0.90	0.84
	Average	12.8	16	0.79	0.72
<i>D. solani</i> IPO 2222, IFB0123		12	16	0.77	0.87
Mean value for all 16 tested <i>D. solani</i> strains		13.9	14.4	0.80	0.77
<i>D. dadantii</i> NCPPB898 ^T IFB0010		12	13	0.79	0.79
<i>D. dianthicola</i> NCPPB453 ^T IFB0103		14	12	0.85	0.91

Phenotypes of strains from different *Dickeya* species

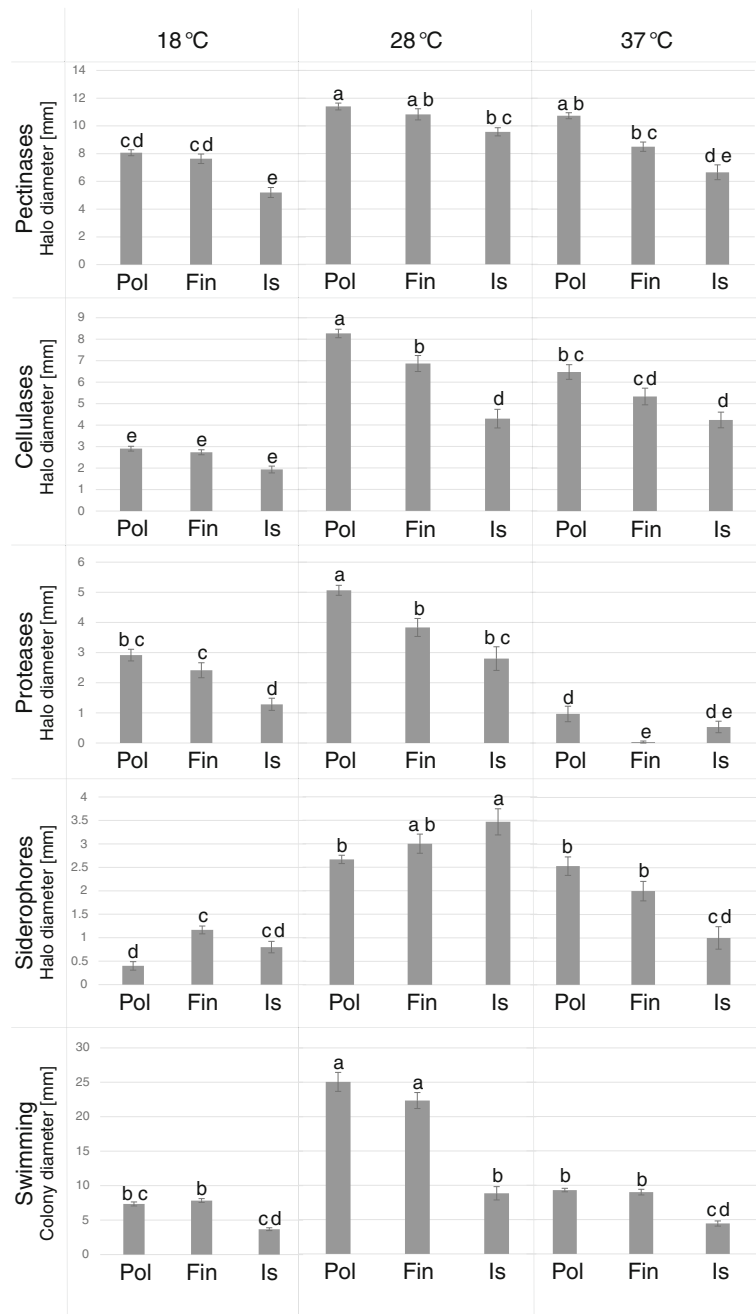
The two strains of *D. dadantii* and three strains of *D. dianthicola* were compared with the *D. solani*^{TS} chosen arbitrarily as a representative. In general, all the strains irrespective of their species have highest pectinolytic, cellulolytic and proteolytic activities at 28 °C (Fig. 3). At this temperature *D. dadantii* and *D. solani* have similar pectinolytic and cellulolytic activities. *D. dianthicola* has significantly lower pectinolytic and cellulolytic activities than two other species. However, at 37 °C *D. dadantii* has significantly higher pectinolytic activity than *D. solani* and *D. dianthicola* (in the case of *D. dianthicola* the activity is close to 0), while at 18 °C *D. solani* possesses the highest pectinolytic activity (similar to the one of *D. dadantii* at 37 °C). *D. solani* indicated the highest proteolytic

activity at 18 and 28 °C (Fig. 3). Concerning the production of siderophores and the swimming ability, no significant differences were observed between the temperature of incubation and the strains tested except of the significantly higher siderophores production by *D. dadantii* strains at 37 °C (Fig. 3).

Total pectinolytic activity of *Dickeya* strains in induced and noninduced conditions

Conditions in which *D. solani* is grown has a significant impact on the total pectinolytic activity especially when evaluated by monitoring spectrophotometrically the formation of unsaturated products from polygalacturonate. In induced conditions (media supplemented with PGA) at 28 °C, all strains of *D. solani* have significantly higher (5

Fig. 2 Comparison of the plant cell wall degrading enzymes activities of *D. solani* strains originating from different countries at 18, 28 and 30 °C. Pol – mean value for five Polish *D. solani* strains: IFB0099, IFB0100, IFB0158, IFB0167, IFB0212; Fin – mean value of halo diameter for five Finnish strains: IFB0231, IFB0236, IFB0254, IFB0261, IFB0265; Is – mean value of halo diameter for five Israeli strains: IFB0124, IFB0125, IFB0455, IFB0456, IFB0457. Error bars represent standard error from three experimental repetitions. Columns with different letters are significantly different when two-way ANOVA followed by post hoc Tukey test at $p < 0.001$ with the F values indicated as follows: pectinases $F = 3.927$, $n = 2$; cellulases $F = 6.717$, $n = 2$; proteases $F = 5.422$, $n = 2$; siderophores $F = 14.981$, $n = 2$; swimming $F = 24.62$, $n = 2$

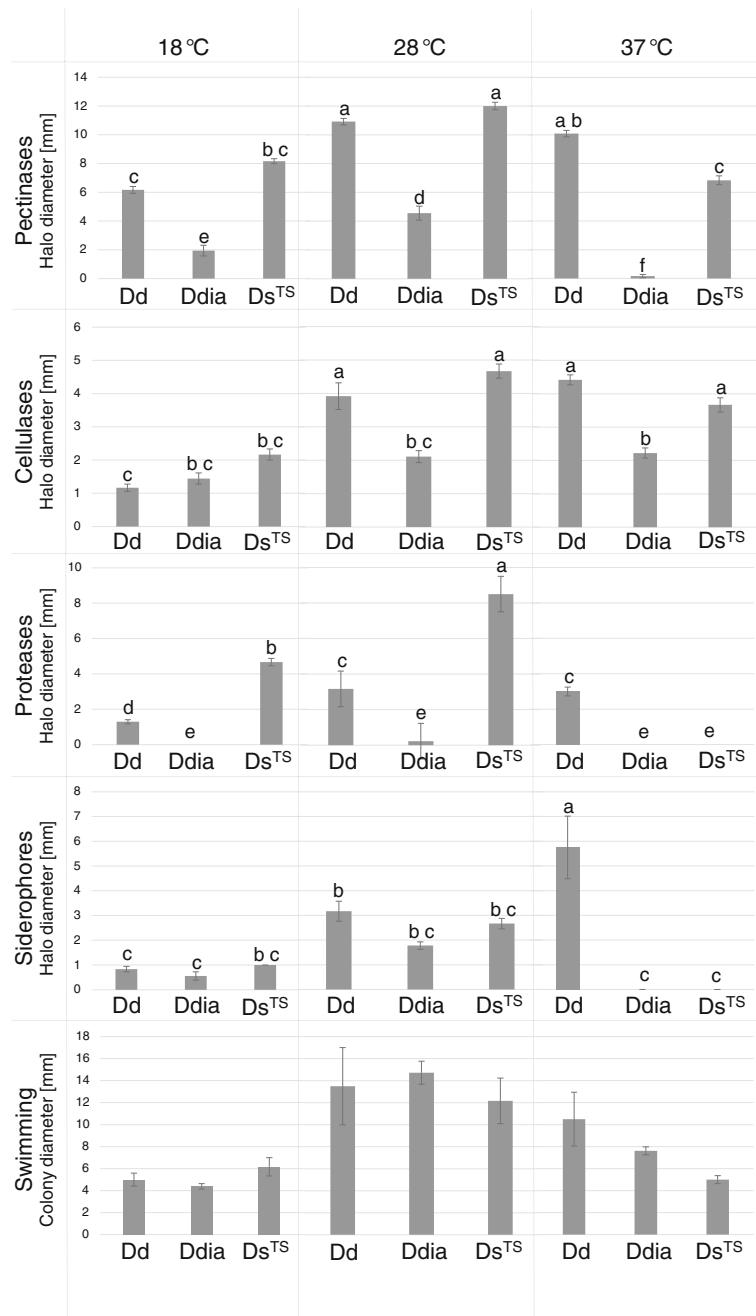


to 10 times) pectinolytic activity than in non-induced conditions (Fig. 4a, Supplement 2). In non-induced conditions the Polish strains have higher pectinolytic activity than the Finnish and the Israeli strains. Also, in induced conditions strains originating from different countries vary significantly from each other. Also in the induced conditions, Polish strains exhibit the highest

pectinolytic activity, while the value for Finnish strains fell between the activities determined for Polish and Israeli strains.

The presence of PGA in the incubation medium also influenced the pectinolytic activity of strains from *D. dadantii* species. *D. dadantii* and *D. solani* strains have significantly higher pectinolytic activity than *D. dianthicola* in both non-induced and induced

Fig. 3 Plant cell wall degrading enzymes activities, siderophore production and swimming motility of different *Dickeya* spp. at 18, 28 and 30 °C. Dd – *D. dadantii*, mean value of halo diameter for two strains: IFB0010 and IFB0016, Ddia – *D. dianthicola*, mean value of halo diameter for three strains: IFB0103, IFB0157 and IFB0188, Ds^{TS} – *D. solani* IPO2222 (IFB0123). Error bars represent standard error from three experimental repetitions. Columns with different letters are significantly different when two-way ANOVA followed by post hoc Tukey test at $p < 0.001$ with the F values indicated as follows: pectinases $F = 23.488$, $n = 2$; cellulases $F = 13.623$, $n = 2$; proteases $F = 221.045$, $n = 2$; siderophores $F = 14.077$, $n = 2$. For swimming the statistical analysis did not distinguish any significant differences ($p > 0.001$, $n = 2$)



conditions (Fig. 4 B). *D. solani* indicated higher pectinolytic activity than *D. dadantii*.

Ability to macerate potato tissue by *D. Solani* from different climatic conditions

Potato slices were inoculated with the bacterial suspensions containing 10^5 , 10^6 and 10^7 cfu ml⁻¹ and

incubated at 28 °C for 48 h. Some variability in the ability to macerate potato tissue was observed among Finish and Israeli strains but not the Polish ones (Supplement 3). The analysis did not distinguish the significance of the inoculum level along with the origin of the *D. solani* strains. However, Polish *D. solani* strains are able to macerate potato tissue more efficiently than the strains from

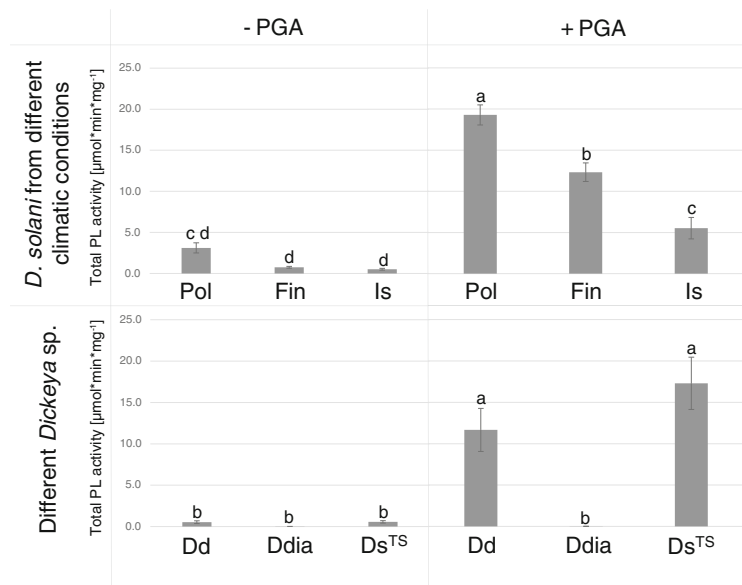


Fig. 4 Comparison of the total pectinolytic activity of tested *Dickeya* strains. **a** *D. solani* strains of different origin in non-induced (–PGA) and induced (+PGA) conditions. **b** strains from different *Dickeya* species. Dd – *D. dadantii*, mean value of halo diameter for two strains: IFB0010 and IFB; Ddia – *D. dianthicola*, mean value of halo diameter for three strains: IFB0103, IFB0157 and IFB0188; Ds^{TS} – *D. solani* IPO2222 (IFB0123); Pol – mean value of halo diameter for five Polish *D. solani* strains: IFB0099, IFB0100, IFB0158, IFB0167, IFB0212; Fin – mean value of halo

diameter for five Finnish strains: IFB0231, IFB0236, IFB0254, IFB0261, IFB0265; Is – mean value of halo diameter for five Israeli strains: IFB0124, IFB0125, IFB0455, IFB0456, IFB0457. Error bars represent standard error from three experimental repetitions. Columns with different letters are significantly different when two-way ANOVA followed by post hoc Tukey test at $p < 0.001$ with the F values indicated as follows: upper panel $F = 7.833$, $n = 3$; lower panel $F = 19.432$, $n = 3$

Finland or Israel at all tested inoculum levels (Fig. 5, Supplement 3). We did not observe any rotting on slices inoculated with the negative

control, so we excluded the possibility of presence of a latent pectinolytic bacteria other than tested inocula.

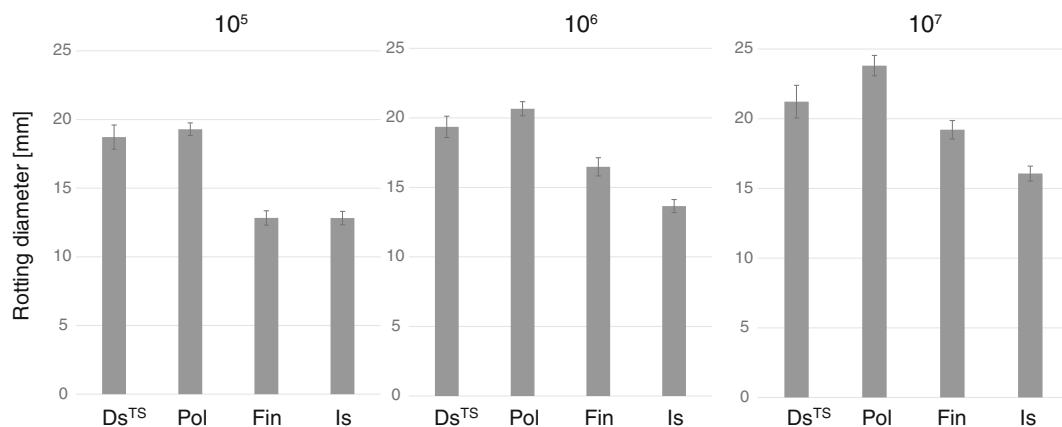


Fig. 5 Comparison of the maceration of potato tuber tissue by *D. solani* strains isolated in different climatic conditions. Ds^{TS} – *D. solani* Type Strain IPO2222; Pol – mean value of halo diameter for five Polish *D. solani* strains: IFB0099, IFB0100, IFB0158, IFB0167, IFB0212; Fin – mean value of halo diameter for five Finnish strains: IFB0231, IFB0236, IFB0254, IFB0261, IFB0265;

Is – mean value of halo diameter for five Israeli strains: IFB0124, IFB0125, IFB0455, IFB0456, IFB0457. Error bars represent standard error from two experimental repetitions. Statistical analysis by two-way ANOVA followed by post hoc Tukey test did not distinguish any significant differences ($p > 0.001$, $n = 9$)

The role of species, origin and temperature on phenotypic traits expression

Performed statistical analysis showed that for enzymatic activities such as pectinases, proteases as well as the ability to chelate iron ions, the species of the strain has higher impact than the temperature of incubation (Table 3a). In the case of pectinases production, the species impact can explain 69 % of the variation. The effect of species also explains 52 % of variability in proteases production and 23 % of variation in the iron chelating ability. On the other hand, the influence of the temperature explains only 15 % of the pectinolytic activity variability and only 1 % of variance both for proteases production and the ability to chelate iron ions. Only for the cellulases activity and the swimming ability the incubation temperature has higher impact on these traits than has the effect of the species, with the temperature explaining 34 % and 32 % of described variability, respectively.

As regards the geographical origin of the strains (country from which *D. solani* strains were isolated), the temperature had higher impact on the exhibited traits (pectinolytic - 30 %, cellulolytic - 41 %, proteolytic - 47 % activities as well as the swimming motility - 47 % and the ability to chelate iron ions - 44 %) than the origin itself (Table 3b). The effect of the climatic origin

of the strains on total pectinolytic activity in induced conditions explains 30 % of the variability. Concerning the ability to macerate potato tissue, the origin of the strains played a more important role than the inoculum level (21 % and 1 %, respectively).

Discussion

The emergence of new, virulent *Dickeya* strains in Europe has become a big problem for the producers of seed potatoes as well as for the farmers producing the ware crop. At the same time, there are no methods of curing the disease and all protection is based on prevention methods, especially important is planting the pathogen-free potato seeds. Of similar significance is testing seed potatoes for the presence of soft rotting bacteria before selling and exporting the seed lots. In the previous years, because of this reason, a zero tolerance policy was introduced in Scotland for *Dickeya* spp. on potato crops (Toth et al. 2011). In Israel, the recent disease outbreaks observed on potatoes grown from tubers imported from the European countries are of great concern, thus their causative agent, *D. solani*, is considered as a quarantine organism in this country (Tsrur (Lahkim) et al. 2013).

Still little is known about the biology, virulence mechanisms and epidemiology of *D. solani*. Genomic analysis of *D. solani* has stated that this species is most closely related to *D. dadantii* 3937 strain (94 % ANI), (Garlant et al. 2013; Pedron et al. 2014; van der Wolf et al. 2014). This is fortunate, because this strain of *D. dadantii* appears to be the best studied in the terms of molecular determinants of virulence and its regulators. A multitude of molecular techniques have been developed to study *D. dadantii* 3937. Potrykus et al. (2014) adapted the techniques used for *D. dadantii* 3937 mutagenesis and applied them in the research on *D. solani*. The major transcriptional regulators of *D. dadantii* 3937 KdgR, PecS, and PecT play a similar role in the regulation of virulence in *D. solani*; they act as the negative regulators of several genes encoding factors involved in the virulence (Potrykus et al. 2014). This study indicates also that the thermo-regulator PecT is an essential regulator of *D. solani* virulence.

The present study of *D. solani* strains originating from different climatic conditions indicated that all of them represent identical rep-PCR and RFLP-PFGE

Table 3 The importance of the species, the geographical origin of the strain and the temperature of incubation on the ability to produce different PCWDE (pectinases, cellulases and proteases) and ability to produce siderophores and swimming expressed in percent's. A The importance of the species and temperature. B. The importance of the geographical origin and the temperature

A		
Factors	Species	Temperature
Pectinases	69	15
Cellulases	25	34
Proteases	52	1
Siderophores	23	1
Swimming	0	32
B		
Factors	Geographical origin	Temperature
Pectinases	20	30
Cellulases	14	41
Proteases	1	47
Siderophores	0,1	44
Swimming	20	47

profiles. This finding confirms the results of Degefu et al. (2013) and van der Wolf et al. (2014), that also indicated high similarities between *D. solani* strains of different origin. The variable tandem repeats method applied by Parkinson et al. (2015) has revealed low variability among 54 tested *D. solani* strains. What is more, 4 out of 16 *D. solani* strains tested in this study (*D. solani*^{TS} IPO2222, Polish IFB0099, Israeli IFB0124 and Israeli IFB0125) have been analyzed by Parkinson et al. (2015). They discovered that *D. solani*^{TS} has a different VNTR profile than three others from the above mentioned strains. However, the observed difference is present only in one locus out of the five tested. Parkinson et al. (2015) stated that the limited variation in VNTR profiles of *D. solani* may reflect a limited number of *D. solani* introductions and/or the short time period since the pathogen's first emergence in Europe. In the same study, 19 VNTR profiles of 45 *D. dianthicola* strains have been distinguished which means that *D. dianthicola* is much more variable than *D. solani* (Parkinson et al. 2015).

In the presented work we tried to answer the question whether observed differences in phenotypic features can be explained by the influence of geographical origin (Poland, Finland and Israel) or the growth temperature (18, 28 and 37 °C). In Poland and Finland, climate is temperate and rather cold (according to Koeppen described in Pidwirny 2006) and it only differs in regard to the summer, which in Poland is temperate and in Finland is shorter and colder. How does it relate to the hot and semi-arid climate in Israel with hot summer? Works of Degefu et al. (2013) and Potrykus et al. (2016) stated that there is an influence of climate change in terms of *D. solani* outbreaks in Finland and Poland. They suggested that the prevalence of *Dickeya* spp. is higher when the summer is hotter and dry. The climate change in the years 1976–2006 shows that the mean temperature values increased in the range of 0.4 to 1.0 °C per decade in Poland and Finland, respectively. The prediction of annual temperature increase in Europe is between 0.3 and 0.35 °C for Poland and Finland respectively, as reported by the European Environmental Agency in 2015 (<http://www.eea.europa.eu/data-and-maps/figures/decadal-average-trends-in-mean-5>).

According to our results *D. solani* strains indicate phenotypic variability apart from their genotypic homogeneity. The analysis of several phenotypic features of fifteen *D. solani* strains originating from three countries with different climatic conditions - Poland, Finland and

Israel - indicated differences in the PCWDE production, the swimming ability and in their ability to macerate potato tissue. Among the tested strains, Polish isolates indicated the lowest phenotypic variability (Supplementary materials 1, 2, 3) and the highest pectinolytic, cellulolytic and proteolytic activities at 18, 28 and 37 °C in addition to the highest ability to macerate potato tissue regardless of the inoculum level. Tsrör (Lahkim) et al. (2013) observed the opposite effect. They stated that *D. solani* strains isolated in Israel are more virulent than those isolated in Europe. It can be explained by the fact that Tsrör (Lahkim) et al. (2013) tested 22 Israeli strains of *D. solani*, we analyzed only five and two of these were isolated from seed potatoes imported to Israel. On the other hand, Tsrör (Lahkim) et al. (2013) used only one strain isolated in Poland, namely IFB0099 (IPO2276), and a few strains isolated in the Netherlands, but none originating from Finland.

Polish strains also had the highest total pectate lyases activity estimated spectrophotometrically. In our studies four out of five Israeli strains exhibited low pectate lyase activity in the applied conditions. On the other hand, the work of Tsrör (Lahkim) et al. (2013) indicated that the strains isolated in Israel had the highest expression of pectate lyase. Observable difference between these reports could be explained by the different methodology that was applied for the estimation of the pectate lyases activities. First of all, we estimated in this study the sum of the activities of several pectate lyases (PelA, PelB, PelC, PelD, PelE, PelL, PelN, PelX), while Tsrör evaluated only the expression of *pelL* gene. Secondly, in the presented work bacteria were grown on M63 Y minimal medium supplemented with PGA as an inducer of the expression of genes encoding pectinolytic enzymes, whereas in the work of Tsrör (Lahkim) et al. (2013) the expression of *pelL* gene was estimated for the bacteria grown in the macerated potato tubers. Former studies performed on *D. dadantii* 3937 showed that different pectate lyases are unequally induced by PGA and other components of the plant tissue (Jafra et al. 1999; Hugouvieux-Cotte-Pattat et al. 2014). Pectate lyases are divided into main and secondary pectate lyases and they are differently regulated. For example the expression of genes encoding the main pectate lyases PelD and PelE are principally induced by PGA, while the expression of genes encoding the secondary pectate lyase PelL is induced by both, PGA and the plant extract (Jafra et al. 1999, Golanowska, not published).

Our studies indicated that the strains' geographical origin (different climates) was less important than the

temperature of incubation, when taking into account the impact of these two variables on the activities of PCWDE. The data presented here reveal significant impact of the genus of bacteria on the level of expression of the genes encoding for pectinases and proteases; *D. solani* strains exhibited higher PCWDE activities and better abilities to cause disease symptoms than *D. dianthicola*. This could explain the phenomenon of *D. solani* having highest chance to cause disease outbreaks when present in the ecological niche together with the other strains from *Dickeya* sp. However, our study includes only 3 strains of *D. dianthicola*, species that is known for high variability.

D. solani strains have higher pectinolytic, cellulolytic and proteolytic activities than the strains of *D. dianthicola* at almost all the tested temperatures and the highest total pectate lyases activities at 28 °C. This could explain the fact that the strains of *D. solani* are better fitted to induce disease symptoms than *D. dianthicola*. Czajkowski et al. (2013) compared *D. solani* to *D. dianthicola* strains and also showed that *D. solani* possesses features which allow for more efficient potato plant colonization than in the case of *D. dianthicola*, especially at 28 °C. This suggests that *D. solani* could be a stronger competitor in the potato ecosystem. In the greenhouse experiments performed at 28 °C, potato roots were more rapidly colonized by *D. solani* than by *D. dianthicola* and 30 days after inoculation higher densities of *D. solani* were found in the stolons and in the progeny tubers. In tubers co-inoculated by vacuum infiltration with GFP-tagged *D. solani* and DsRed-tagged *D. dianthicola*, *D. solani* strain outcompeted *D. dianthicola* (Czajkowski et al. 2010). Moreover, *D. solani* indicated susceptibility only to 31 % of saprophytic bacteria, potential biocontrol agents, when *D. dianthicola* indicated susceptibility to 80 % of the tested strains (the selection included strains from the species such as: *Serratia plymuthica*, *Pseudomonas* spp., *Delftia acidovorans*, *Pantoea agglomerans*, *Delftia acidovorans*, *Rahnella* sp., *Klebsiella* sp. and *Acinetobacter* sp. (Czajkowski et al. 2010)).

In conclusion, our study showed that *D. solani* strains are genetically homogenous, but they exhibit phenotypic differences. Moreover, *D. solani* strains have higher plant cell-wall degrading enzymes activities (such as pectinases, cellulases and proteases) than *D. dianthicola* strains, regardless of the temperature of incubation. Besides, Polish *D. solani* strains are superior to strains

originating from Finland and Israel in the production of PCWDE and their ability to macerate potato tissue.

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